EXHIBIT A:

MOUSE LKB1 COMPARED TO HUMAN LKB1

BLAST

Basic Local Alignment Search Tool

Edit and Resubmit Save Search Strategies Formatting options Download

Blast 2 sequences

SEQ ID NO: 6 (44-343) Compared to Mouse LKB1

Results for: IcI|41549 SEQ ID NO: 6(433aa)

Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

Query ID Ici|41549 Ici|41549

Description SEQ ID NO: 6

Molecule type amino acid Query Length

Subject ID

gi[7106425[ref]NP_035622.1]

ripidon serinathroonine-protein kinase 11 [Mus musculus] >qi81917862(sp)Q8WTK7.1|STX11_MOUSE RenName: Full=Serinathroonine-protein kinase 11 [Mus musculus] >qi81917862(sp)Q8WTK7.1|STX11_MOUSE RenName: Full=Serinathroonine-protein kinase LK81 [Mus musculus] >qi8194679(sp)para syndrome kinase LK81 [Mus musculus] >qi8194679(sp)para syndrome kinase LK81 [Mus musculus] >qi8194679(sp)qaba22700.1] serinathroonine-protein kinase LK81 [Mus musculus] >qi819469(sp)qab22700.1] serinathroonine-protein kinase LK81 [Mus musculus] >qi819469(sp)qab22700.1] serinathroonine-protein kinase LK81 [Mus musculus] >qi79469(sp)qab22700.1] serinathroonine-lakane 11 [Mus musculus] >qi79469(sp)qab22700.1] serinathroonine-lakane 11 [Mus musculus] >qi7947(sp)qab22700)[Mus musculus] >qi7947(sp)qab22700][Mus musculus] >qi7947(sp)qab2270][Mus musculus] >qi7947(sp)qab2270][Mus musculus] >qi7947(sp)qab2270[Musculus] >qi7947(sp)qab227

Molecule type amino acid

Subject Length

436

BLASTP 2.2.24+ Citation

Reference Stephen F, Allschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghul Zhang, Zhang Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402

and Professional state glaterature of professional state and state of the Referency - compositional score matrix addicational Stephen F. Altschut, John C. Wootton, E. Michael Gentz, Richa Agarwala, Aleksandr Mongulis, Alejandro A. Schäffer, and Yi-Kuo Yu (2005) "Protein databases searches using compositionally adjusted substitution matrices", FEBS J. 272:5101-5109.

Other reports: Search Summary [Taxonomy reports] [Multiple alignment]
Search Parameters

Search parameter name Search parameter value

Query range	44-343		
Word size	3		
Expect value	10		
Hitlist size	100		
Gapcosts	11,1		
Matrix	BLOSUM62		
Low Complexity Filter	Yes		
Filter string	L;		
Genetic Code	1		
Window Size	40		
Threshold	11		
Companition beneat state	9		

Karlin Allechul statistics

Params Ungapped Gapped

Lambda	0.320459	0.267	
K	0.139595	0.041	
н	0.429592	0.14	

Results Statistics

Results Statistics parameter name Results Statistics parameter value

Graphic Summary

Distribution of Blast Hits on the Query Sequence

An overview of the database sequences aligned to the query sequence is shown. The score of each alignment is indicated by one of fine different colors, which divides the range of scores into the groups. Multiple silgnments or the semination are consisted by a subject fine. Nuclear, over a hit which divides the range of scores into the groups. Multiple silgnments or the semination over a hit sequence takes the user to the associated alignments. Never This spacific is an invertised of database sequences alignment of the sequence. Alignments are coher-coded by cone, within one of the score ranges. Multiple alignments on the same database sequences are connected by a dashed line. Mousing over an alignment shows the alignment definition and soon in the box of the top. Clicking an elignment disjoint by the alignment database.

Dot Matrix View /

Plot of IcII41549 vs ail7106425|refiNP 035622.1| [?]

This dot matrix view shows regions of similarity based upon the BLAST results. The query sequence is represented on the X-axis and the numbers represent the bases/residues of the quarter than the properties of the properties are shown in the piot as lines. Plus stand and protein matches are altered from the bottom defined in the upper right corner, minus stand matches are shared from the potenties of the properties of the

×

Descriptions

Score Sequences producing significant alignments: (Bits) ref[pp 035622.1] serine/threonine-protein kinase 11 [Mus musc... 526 4e-154

Alianments

>ref|NP 018622.1| serine/threonine-protein kinase 11 [Mas masoulus] sp|08987.1|3781_M0088 RocName: Full-Serine/threonine-protein kinase 11, AltHame: Full-Serine/threonine-protein kinase 11, AltHame: Full-Serine/threonine-protein kinase LR3| cinase LKB1 gb|AAD31044,1|AP145287 1 Peutz-Jeghers syndrome kinase LKB1 [Mus musculus] 9 more sequence titles

sh|AP71370.1|ar151711] protein kinase LKD1 [Mus musculus]
sh|AP71370.1|ar151711] protein kinase LKD1 [Mus musculus]
sh|AD51200.2| lecting Chirconin-protein kinase LXD1 [Mus musculus]
sh|AD51270.1| Fertir-Speches syndrome protein [Mus musculus]
sh|AD51271.1| Sertir-Arecaine kinase 1.1 [Mus musculus]
sh|AB71271.1| Sertir-Arecaine kinase 1.1 [Mus musculus]
sh|AB74201.1| Sertir-Arecaine kinase 1.1 [Mus musculus]
sh|AB74201.1| Sertir-Arecaine kinase 1.1 [Mus musculus]
sh|AB74201.1| Sertir-Arecaine kinase 1.1 [mus musculus]

Score = 525 bits (1354), Expect = 4e-154, Method: Compositional matrix adjust Identities = 268/300 (96%), Positives = 292/300 (98%), Gaps = 0/300 (0%)

KLIGKYIMGDLLGBGSYGKVKEVLDSETLCRRAVKilkKkklrriPNGEANVKKEIQLLR KLIGKYIMGDLLGBGSYGKVKEVLDSETLCRRAVKILKKKKLRRIPNGEANVKKEIQLLR KLIGKYLMGDLLGEGSYGKVKEVLDSETLCRRAVKILKKKKLRRIPNGEANVKKEIQLLR 103 RLEHANVIQIADVLYMEEKQKMYMVMEYCVCGMQEMLDGVPEKRPPVCQAHGYPCQLIDG RLEH-HVIQLVDVLYMEEKQKMYMVMRYCVCGMQEMLDSVPEKRPPVCQAHGYP QLIDG RLEHRNVIQLVDVLYMEEKQKMYMVMSYCVCGMQEMLDSVPEKRPPVCQAHGYPCLIDG 104 Ouerv 163 LEYLHSGGYVHKDIKPgnillttggtlKISDLGVARALHFFAADDTCKTSGGSPAFQPPE LEYLHSGGYVHKDIKPGNLLLTT GTLKISDLGVARALHFFA DDTCKTSGGSPAFQPPE LEYLHSGGYVHKDIKPGNLLLTTNGTLKISDLGVARALHFFAVDDTCKTSGGSPAFQPPE Query 164 Sbjet 164 langldtfsgfrydiksagytlynittglypfsgdniyklpenigkgsyaifgdcgfpls iangldtfsgfrydiksagytlynittglypfsgniyklfenig4g + if Dcgfpls iangldtfsgfrydiksagytlynittglypfsgniyklpenigagdfilqdcgfls 283 Sbjct 224 Query 284 DILKOMLEYEPAKRPSIRQIRQHSHFRKKHDPGCODVDIDPSDTKDRNRSMTVVFYLED SDJC 284 DILKOMLEYEPAKRPSIRQIRQHSHFRKKHD AEA VDIPPSDTKDRNRSMTVVFYLED DILKOMLEYEPAKRSIRQIRQHSHFRKHDLEALVEPTPSDUTTDRNRSMTVVFYLED